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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
10/591,521	09/01/2006	Hiroharu Kawahara	125192.00501	1684
Pepper Hamilto	7590 12/24/200 n	EXAMINER		
500 Grant Stree Pittsburgh, PA	t, 50th Floor	KIM, ALEXANDER D		
riusburgh, rA	13219		ART UNIT	PAPER NUMBER
			1656	
			MAIL DATE	DELIVERY MODE
			12/24/2008	PAPER

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

Advisory Action Before the Filing of an Appeal Brief

Application No.	Applicant(s)		
10/591,521	KAWAHARA, HIROHARU		
Examiner	Art Unit		
ALEXANDER D. KIM	1656		

	ALEXANDER D. KIM	1656	
The MAILING DATE of this communication appear	ars on the cover sheet with the c	orrespondence add	ress
THE REPLY FILED <u>03 December 2008</u> FAILS TO PLACE THIS	APPLICATION IN CONDITION FO	OR ALLOWANCE.	
1. The reply was filed after a final rejection, but prior to or on application, applicant must timely file one of the following rapplication in condition for allowance; (2) a Notice of Appe for Continued Examination (RCE) in compliance with 37 C periods:	eplies: (1) an amendment, affidavit al (with appeal fee) in compliance v	, or other evidence, w with 37 CFR 41.31; or	hich places the (3) a Request
a) The period for reply expiresmonths from the mailing b) The period for reply expires on: (1) the mailing date of this Ac no event, however, will the statutory period for reply expire la Examiner Note: If box 1 is checked, check either box 700 or (1)	dvisory Action, or (2) the date set forth interthan SIX MONTHS from the mailing	date of the final rejection	n.
MONTHS OF THE FINAL REJECTION. See MPEP 706.07(f Extensions of time may be obtained under 37 CFR 1.136(a). The date of have been filed is the date for purposes of determining the period of extender 37 CFR 1.17(a) is calculated from: (1) the expiration date of the slipset forth in (b) above, if checked. Any reply received by the Office later may reduce any earned patent term adjustment. See 37 CFR 1.704(b). NOTICE OF APPEAL	n which the petition under 37 CFR 1.13 ension and the corresponding amount on the reply origin	of the fee. The appropria nally set in the final Offic	te extension fee e action; or (2) as
 The Notice of Appeal was filed on A brief in compl filing the Notice of Appeal (37 CFR 41.37(a)), or any exten Notice of Appeal has been filed, any reply must be filed with 	sion thereof (37 CFR 41.37(e)), to	avoid dismissal of the	
AMENDMENTS	a dente de las reelles bolance	=201 a (b a (a a) b	
3. The proposed amendment(s) filed after a final rejection, b (a) They raise new issues that would require further con (b) They raise the issue of new matter (see NOTE below	sideration and/or search (see NOT v);	E below);	
(c) ☐ They are not deemed to place the application in bett appeal; and/or	er form for appeal by materially rec	lucing or simplifying tr	ie issues for
(d) ☐ They present additional claims without canceling a c NOTE: (See 37 CFR 1.116 and 41.33(a)).	orresponding number of finally reje	cted claims.	
4. 🔲 The amendments are not in compliance with 37 CFR 1.12	 See attached Notice of Non-Cor 	mpliant Amendment (F	PTOL-324).
5. Applicant's reply has overcome the following rejection(s):			
6. Newly proposed or amended claim(s) would be allow non-allowable claim(s).	·	•	_
7. For purposes of appeal, the proposed amendment(s): a) [how the new or amended claims would be rejected is prov The status of the claim(s) is (or will be) as follows: Claim(s) allowed:		be entered and an ex	planation of
Claim(s) objected to: Claim(s) rejected: <u>20 and 21</u> . Claim(s) withdrawn from consideration:			
AFFIDAVIT OR OTHER EVIDENCE			
 The affidavit or other evidence filed after a final action, but because applicant failed to provide a showing of good and was not earlier presented. See 37 CFR 1.116(e). 			
 The affidavit or other evidence filed after the date of filing a entered because the affidavit or other evidence failed to over showing a good and sufficient reasons why it is necessary 	rercome <u>all</u> rejections under appear and was not earlier presented. Se	l and/or appellant fails e 37 CFR 41.33(d)(1)	s to provide a
 The affidavit or other evidence is entered. An explanation REQUEST FOR RECONSIDERATION/OTHER 	of the status of the claims after er	try is below or attache	ed.
 The request for reconsideration has been considered but <u>See Continuation Sheet.</u> 		condition for allowand	ce because:
12. ☐ Note the attached Information <i>Disclosure Statement</i>(s). (l13. ☐ Other:	PTO/SB/08) Paper No(s)		
	/Rebecca E. Prouty/ Primary Examiner, Art U	nit 1652	

Continuation of 11. does NOT place the application in condition for allowance because: Applicant's amendment after final rejection, filed on 12/09/2008, is acknowledged and has been entered. The said amendment does not appear to have any changes from the previous claims before Final Office Action.

Previous objection to the specification is withdrawn by the virtue of Applicants' amendment in the specification reciting "SC-02MFP" and "SC-01MFP".

Applicants' arguments in the amendment filed on 12/09/2008 have been fully considered. However, applicant's arguments are not found persuasive to overcome the outstanding rejection(s) as set forth in the Final Office action mailed on 10/03/2008 for the reasons of record stated therein. Applicants argue that the instant specification (paragraph 0026-0033) on pages 6-8 specifically describe how SC-02MFP may be produced form RMP18226; and SC-01MFP may be produced form KMS-12BM. The instant specification does not used paragraph numbers in the newly filed specification (filed on 12/3/2008); thus, entire pages 6-8 have been considered.

The specification disclose the cell strain SC-01MFP were established from the RPMI8226 cell strain of human myeloma origin; and the SC-02-MFP were established from FERM BP-10077 (see bottom of page 8). However, it is still not clear how to decipher the method of making SC-01MFP and SC-02-MFP such that they are structurally different from the original cell RPM18226 and FERM BP-10077, respectively. The specification, bottom of page 6 to top of page 7, recites that "the inventors of this invention acquired a mutated strain by isolating and selecting a human cell strain form various kinds of human sources that allows a long term stable protein production"; wherein the selection encompasses selecting by the total weight of intracellular protein of cells (see page 7, lines 11-16) and also selecting for cloning rate and doubling time (see page 7, lines 18-24). It is unclear how said isolating and selecting step would produce a mutant strain.

The specification states later on page 8 that "The variant cells obtained as described were induced to mutate in a medium in which nitrosoguanidine, a carcinogenic substance, had been added" and "selected" the mutant. However, it is unclear what kinds of genetic change(s) [i.e., a mutant] were made by the presence of nitrosoguanidine since some population may undergo mutation (if any, since the concentration is unknown and mutagen induced mutation which is concentration dependent) while some population may not undergo mutation. In view of instant specification, there is no evidence that SC-01MFP and SC-02-MFP are mutants induced by the presence of nitrosoguanidine derived from the original cell RPM18226 and FERM BP-10077, respectively. Also, the selection step of cell having cloning rate of over 90% do not contribute to making a mutant as described earlier. Also, the recitation of "The variant cells" in the beginning of page 8 is unclear if it is referring to the various original human cell strain described on top of page 7; or it is referring to the cells after the selection and cloning as described on bottom of page 7 which already selected for over 90% cloning rate.

The specification on page 8, bottom, indicates that the criteria of continuous protein expression of certain period is the criteria for establishing as human cell strain for protein production. However, a cell will continuously expresses a protein as long as a cell is kept in a viable culture condition and the culturing the cell would not contribute to any structural changes such as forming a mutant. As noted in the previous office action, since the Office does not have the facilities for examining and comparing applicants' claimed cell with the cells of the prior art, the burden is on applicant to show a novel or unobvious difference between the claimed product and the product of the prior art (i.e., that the protein of the prior art does not possess the same material structural and functional characteristics of the claimed protein). See In re Best, 562 F.2d 1252, 195 USPQ 430 (CCPA 1977) and In re Fitzgerald et al., 205 USPQ 594. Thus, Claims 20 and 21 are remain rejected over Pene et al. and Hata et al., respectively..